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INTERNATIONAL	Inter 11	auonai Bureau .	
INTERNATIONAL APPLICATION PUBL (51) International Patent Classification 6:	JSHED I	UNDER THE PATENT COOPERATION	ON TREATY (PCT)
C11D 3/39	A1	(11) International Publication Number:	WO 95/13352
		(43) International Publication Date:	18 May 1995 (18.05.95
(22) International Filing Date: (30) Priority Data: (31) 08/151717 (32) 12 November 1993 (12.1) (33) 278664 (34) 21 July 1994 (21.07.94) (35) Applicant (for all designated States except AU BB LK MN MW NZ SD): UNILEVER N.V. [NL/N 455, NL-3013 AL Rotterdam (NL). (37) Applicant (for AU BB CA GB IE LK MN MW NZ UNILEVER PLC [GB/GB]: unilever House, I London EC4 4BQ (GB). (38) Priority Data: (48) 1996 (GB/GB): UNILEVER N.V. [NL/N MW NZ UNILEVER PLC [GB/GB]: unilever House, I London EC4 4BQ (GB). (57) Inventors: MADISON, Stephen, Alan; 24 Oriole F City, NY 10956 (US). COOPE, Janet, Lynn; 24 Avenue, Cliffside Park, NJ 07010 (US).	1.93) U. CA GB III NLJ; Weens Z SD only): Blackfriars. Road, New 20 Lawton	CN, CZ, DE, DK, ES, FI, GB, KR, KZ, LK, LT, LV, MD, MC PL, PT, RO, RU, SD, SE, SI, European patent (AT, BE, CH, IE, IT, LU, MC, NL, PT, SE), CG, CI, CM, GA, GN, ML, MR, patent (KE, MW, SD, SZ). Published With international search report. Before the expiration of the tin claims and to be republished in amendments.	BB, BG, BR, BY, CA. CH GE, HU, IP, KE, KG, KP G, MN, MW, NL. NO. NZ SK, TJ. TT, UA, UZ, VN. DE, DK, ES, FR, GB, GR OAPI patent (BF, BJ, CF, NE, SN, TD, TG), ARIPO
(4) Title: IMINE QUATERNARY SALTS AS BLEACE (7) Abstract	H CATAL	YSTS	
Novel bleach catalysts, a method for bleaching subset reported. The bleaches are quaternary imine salts. Sublis and a peroxygen compound.	strates usin ostrates suc	g these catalysts and detergent compositions h as fabrics may be bleached in an aqueous s	containing the catalysts solution containing these

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IMINE QUATERNARY SALTS AS BLEACH CATALYSTS BACKGROUND OF THE INVENTION

Field of the Invention

5 The invention relates to a new type of low-temperature bleaching system and a method of cleaning substrates therewith.

The Related Art

- Many household and personal care products are formulated with an active oxygen-releasing material to effect removal of stain and soil. Oxygen-releasing materials have an important limitation; their activity is extremely temperature-dependent. Temperatures in excess of 60°C are
- normally required to achieve any bleach effectiveness in an aqueous wash system. Especially for cleaning fabrics, high temperature operation is both economically and practically disadvantageous.
- The art has partially solved the aforementioned problem through the use of activators. These activators, also known as bleach precursors, often appear in the form of carboxylic acid esters. In an aqueous liquor, anions of hydrogen peroxide react with the ester to generate the
- corresponding peroxyacid which oxidizes the stained substrate. Commercial application of this technology is found in certain fabric bleaching detergent powders incorporating tetraacetylethylenediamine (TAED) and sodium nonanoyloxybenzene sulfonate (SNOBS).

TAED is effective only under warm-hot wash conditions, i.e. above 30°C. Although this material is widely employed in Europe with laundry detergent, cold water consumer washing habits have not permitted use in the United States. SNOBS

can operate at lower temperatures than TAED. For this reason, it has been commercialized in the United States but its performance could still be improved.

Another problem with carboxylic acid ester precursors such as TAED and SNOBS is that conversion to peracid is inefficient. A further difficulty is that they are not catalytic. Once the ester has been perhydrolyzed, it can no longer be recycled. Accordingly, relatively large amounts of precursor are necessary. Amounts as high as 8% may be necessary in a detergent formulation for bleaching fabrics. At such high use levels, cost for these relatively expensive chemicals is of major concern.

10

Recently there has been reported in U.S. Patent 5,047,163, U.S. Patent 5,045,233 and U.S. Patent 5,041,232, all to Batal and Madison, a system for activating bleach precursors based upon sulfonimines and N-

- sulfonyloxaziridines. While these compounds have been shown to be highly effective, even better catalysts are sought, especially for wash temperatures around 10°C, such as are experienced in Japan.
- Outside the context of consumer products, there have been reports of catalytic oxidizing agents. Hanquet and coworkers, in a series of articles, reported preparation of a new class of stable olefin epoxidizing agents, namely oxaziridinium salts. See Hanquet, Lusinchi and Milliet,
- 25 Tet, Let. (1988) 3941; Hanquet, Lusinchi and Milliet, C.R. Acad. Sci. Paris (1991) Series II, 625; and Hanquet, Lusinchi and Milliet, Tet, Let. (1988) 2817. These oxaziridinium salts were prepared by peracid or monopersulfate oxidation of a corresponding quaternary
- imine salt under alkaline conditions. Epoxides were reported to be formed from the reaction of olefins with the oxaziridinium salts. Reactions were conducted either in organic solvents or in organic solvent-water biphasic media. Beyond use as a synthetic tool, there is no
- suggestion of any possible application for quaternary imine salt chemistry to the problem of removing stains in consumer applications, such as in cleaning fabrics.

WO 95/13352

It is an object of the present invention to provide an improved bleaching system and detergent composition containing such system that operates over a wide temperature range including that of under 30°C.

5

It is another object of the present invention to provide novel bleach catalysts which are effective at relatively low concentrations, thereby achieving a cost-effective stain removal system.

10

A still further object of the present invention is to provide a method for bleaching stained substrates such as clothes, household hard surfaces including sinks, toilets and the like, and even dentures.

15

Other objects of the present invention will become more readily apparent through the following summary, detailed description and examples.

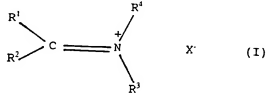
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SUMMARY OF THE INVENTION

A bleaching composition is provided comprising:

- (i) from about 1 to about 60% by weight of a peroxygen compound;
- 25 (ii) from about 0.01 to about 10% of an oxygen transfer agent whose structure is:

30



35

40

wherein:

R' and R' may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl and cycloalkyl radicals;

 ${\ensuremath{\mathsf{R}}}^2$ may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, cyano, alkoxy, keto, carboxylic and carboalkoxy radicals;

 ${\ensuremath{\mathsf{R}}}^3$ may be a substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, and

 \mathbb{R}^1 with \mathbb{R}^2 and \mathbb{R}^2 with \mathbb{R}^3 may respectively together form 10 a radical selected from the group consisting of cycloalkyl, polycyclo, heterocyclic and aromatic ring systems;

X is a counterion stable in the presence of oxidizing agents; and

(iii) from about 0.5 to 50% of a surfactant.

15

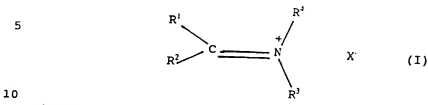
Additionally, there is provided a method for bleaching a stained substrate comprising the step of applying to the stained substrate an aqueous solution comprising a peroxygen compound and an oxygen transfer agent whose 20 structure is $R^1R^2C=N^+R^3R^4X^$ defined above, the mole ratio of peroxygen compound to with radical groups as oxygen transfer agent being from about 250:1 to about 1:2.

DETAILED DESCRIPTION

25

It has been found that certain types of quaternary imine salts can operate as catalysts on peroxygen compounds to transfer active oxygen to stains. Consumer and industrial articles can effectively be bleached to remove stains present on such articles. Thus, quaternary imine salt chemistry is more than a synthetic curiosity as in the conversion of olefins to epoxides reported by Hanquet et al. Unlike the Hanquet et al. reaction medium that requires an organic solvent, quaternary imine salts can be devised 35 for use in completely aqueous wash systems.

Quaternary imine salts covered by the present invention are those whose structure is:



wherein:

30

R¹ and R⁴ may be hydrogen or a C₁-C₃₀ substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl and cycloalkyl radicals;

R² may be hydrogen or a C₁-C₃₀ substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, cyano, alkoxy, keto, carboxylic and carboalkoxy
20 radicals;

 R^3 may be a C_1 - C_{30} substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, and cyano radicals;

25 R¹ with R² and R² with R³ may respectively together form a cycloalkyl, polycyclo, heterocyclic or aromatic ring system;

 X^{\cdot} is a counterion stable in the presence of oxidizing agents.

Heterocyclic rings according to this invention include cycloaliphatic and cycloaromatic-type radicals, incorporating an oxygen, sulfur and/or nitrogen atom within the ring system. Representative nitrogen heterocycles include pyridine, pyrrole, imidazole, triazole, tetrazole, morpholine, pyrrolidine, piperidine and piperazine. Suitable oxygen heterocycles include furan, tetrahydrofuran and dioxane. Sulfur heterocycles may include thiophene and tetrahydrothiophene.

Counterion X may be selected from chloride, bromide, sulfate, methosulfate, sulfonate, p-toluenesulfonate, borontetrafluoride, PF_5 , phosphate and cyano radicals.

5 The term "substituted" is defined in relation to $R^{1},\ R^{2},\ R^{3}$ and R^4 as a substituent which is a nitro, halo, cyano, $C_1 - C_{20}$ alkyl, amino, aminoalkyl, thioalkyl, sulfoalkyl, carboxyester, hydroxy, C_1-C_{20} alkoxy, polyalkoxy or C_1-C_{40} quaternary di- or tri-alkylammonium function. 10

The most preferred quaternary imine salts are the 3,4dihydroisoquinolinium salts of structure II where $\ensuremath{R^5}$ and $\ensuremath{R^0}$ are defined by the same radicals as those for $\ensuremath{\mathsf{R}}^2$:

15

20

Table I lists specific illustrative compounds represented by structure II.

TABLE I

			TABLE	_1_		
	Compou	nd R ⁴	R ⁵		R ⁶ Y	===
	1	CH,	Н			
5	5 2	СН,	Н		BF ₄ -	
	3	СН3	СН,		p-tosyla	te
	4	СН,	NO ₂	F		
	5	СН3	Cl	H		
	6	CH ₃	осн,	H	- DI 4	
10	7	phenyl		Н	DIOSYTATE	€.
	8	benzyl	H phasul	н	CH ₃ SO ₄	
	9	(CH ₂) ₂ OH	phenyl	Н Н		
	10	CH ₃	Ch coon	H	PF ₆	
	11	(CH ₃) ₂ CH	CH2COCH3	H	PF ₆	
15	12	CH,	COCH,	Н	CH ₂ CH ₃ SO ₄	
	13	(CH ₃ (CH ₂) ₁₁	SO ₂ -Na +	Н	CJ.	
	14	(CH ₂ (CH ₂) ₁₅	H	H	p-tosylate	₃.
	15	CH ₂ CH ₂ N (CH ₂),	Br	Н	CH ₂ SO ₄ ·	
	16	CH ₃		H	C1-	
20	17	CH ₃	F	Н	CJ.	
	18	CH ₃	CF ₃	Н Н	PF ₆	
	19	CH ₃	CH ₂ OPO ₃ Na ₂	H	C1.	
	20	2-pyridyl	pyridyl	H	CJ.	
ĺ	21	CH ₃	Н	Н	Cl.	
25	22	CH ₁ CH ₂ O(CH ₂) ₂	CH ₂ N ⁺ (CH ₃) ₃	Н	CH,SO4	7
	23	CH,	H	Н	CH3SO4	
	24	CH ₃	(CH ₂) ₇ CH ₃	Н	Cl-	7
	25	(CH ₂) ₇ CH ₃	CO ₂ Na+	Н	Cl.	1
	26	CH ₃	Н	Н	p-tosylate	1
30	27	CH ₃	H	CH ₃	C1.	1
<u></u>		Cri ₃	Н	phenyl	C1.	
						נ

Additional compounds according to the present invention are outlined below as structures III through χ .

$$C = N$$
 $C = N$
 CH_3
 BF_4
 CH_3

$$C = N CH_3$$

$$CH_3 BF_4$$
(VIII)

$$CH = N$$

$$CI^{-}$$
(X)

The foregoing oxygen transfer agents may be incorporated into detergent bleach compositions along with a further essential component which is a peroxygen compound capable of yielding peroxide anion in an aqueous solution.

Amounts of oxygen transfer agent suitable for the present invention may range from about 0.01 to 10%, preferably from about 0.2 to 5%, optimally from about 0.5 to 1.5% by weight of the composition.

5

The peroxygen compound may be present from about 1 to 60%, preferably from about 1.5 to 25%, optimally from about 2 to 10% by weight.

The molar ratio of peroxide anion (or a peroxygen compound generating the equivalent amount of peroxide anion) to oxygen transfer agent will range from about 1500:1 to about 1:2, preferably from about 150:1 to 1:1, optimally from about 60:1 to 3:1.

15

Peroxide anion sources are well known in the art. They include the alkali metal peroxides, organic peroxides such as urea peroxide, and inorganic persalts, such as the alkali metal perborates, percarbonates, perphosphates, persilicates and persulfates. Mixtures of two or more such compounds may also be suitable. Particularly preferred are sodium perborate tetrahydrate and, especially, sodium perborate monohydrate. Sodium perborate monohydrate is preferred because it has excellent storage stability while also dissolving very quickly in aqueous solutions.

Alkyl hydroperoxides are another suitable class of peroxygen compounds. Examples of these materials include cumene hydroperoxide and t-butyl hydroperoxide.

30

35

Organic peroxyacids may also be suitable as the peroxygen compound. Such materials have a general formula:

wherein R is an alkylene or substituted alkylene group containing from 1 to about 22 carbon atoms or a phenylene or substituted phenylene group, and Y is hydrogen, halogen, alkyl, aryl or

5

The organic peroxyacids usable in the present invention can contain either one or two peroxy groups and can be either aliphatic or aromatic. When the organic peroxyacid is aliphatic, the unsubstituted acid has the general formula:

15
$$0 \text{ HO-O-C-} (CH_2)_n - Y$$

where Y can be, for example, H, CH_3 , CH_2Cl , COOH, or COOOH; and n is an integer from to 1 to 20.

20 When the organic peroxy acid is aromatic, the unsubstituted acid has the general formula:

25 wherein Y is hydrogen, alkyl, haloalkyl, halogen or COOH or COOH.

Typical monoperoxyacids useful herein include alkyl peroxyacids and aryl peroxyacids such as:

- (i) peroxybenzoic acid and ring-substituted peroxybenzoic acid, e.g. peroxy-α-naphthoic acid;
 - (ii) aliphatic, substituted aliphatic and arylalkyl monoperoxy acids, e.g. peroxylauric acid, peroxystearic acid, and N,N-phthaloylaminoperoxycaproic acid (PAP); and
- (iii) amidoperoxyacids, e.g. monononylamide of either peroxysuccinic acid (NAPSA) or of peroxyadipic acid (NAPAA).

Typical diperoxyacids useful herein include alkyl diperoxyacids and aryldiperoxy acids, such as:

- (i) 1,12-diperoxydodecanedioic acid;
- (ii) 1,9-diperoxyazelaic acid;
- (iii) diperoxybrassylic acid; diperoxysebacic acid and 5 diperoxyisophthalic acid;
 - (iv) 2-decyldiperoxybutane-1,4-dioic acid;
 - (V) 4,4'-sulfonylbisperoxybenzoic acid; and
- (vi) N,N'-terephthaloyl-di(6-aminoperoxycaproic acid) 10 (TPCAP).

Particularly preferred organic acids are peracetic acid, monoperoxyphthalic acid (magnesium salt hexahydrate), PAP, TPCAP and diperoxydodecanedioic acid. Under certain 15 circumstances, hydrogen peroxide itself may directly be em: yed as the peroxygen compound.

Bleach systems of the present invention may be employed for a wide variety of purposes, but are especially useful in 20 the cleaning of laundry. When intended for such purpose, the peroxygen compound and oxygen transfer agent of the present invention will usually also be combined with surface-active materials, detergency builders and other known ingredients of laundry detergent formulations.

25

The surface-active material may be naturally derived, such as soap or a synthetic material selected from anionic, nonionic, amphoteric, zwitterionic, cationic actives and mixtures thereof. Many suitable actives are commercially 30 available and are fully described in the literature, for example in "Surface Active Agents and Detergents", Volumes I and II, by Schwartz, Perry and Berch. The total level of the surface-active material may range up to 50% by weight, preferably being from about 1% to about 40% by weight of 35 the composition, most preferably 4 to 25%.

Synthetic anionic surface-actives are usually water-soluble alkali metal salts of organic sulfates and sulfonates having alkyl radicals containing from about 8 to about 22 carbon atoms.

5

Examples of suitable synthetic anionic detergent compounds are sodium and ammonium alkyl sulfates, especially those obtained by sulfating higher (C_g-C_{1g}) alcohols produced, for example, from tallow or coconut oil; sodium and ammonium 10 alkyl (C_9-C_{20}) benzene sulfonates, particularly sodium linear secondary alkyl (C_{10} - C_{15}) benzene sulfonates; sodium alkyl glyceryl ether sulfates, especially those ethers of the higher alcohols derived from tallow or coconut oil and synthetic alcohols derived from petroleum; sodium coconut 15 oil fatty acid monoglyceride sulfates and sulfonates; sodium and ammonium salts of sulfuric acid esters of higher $(C_9 - C_{18})$ fatty alcohol-alkylene oxide, particularly ethylene oxide reaction products; the reaction products of fatty acids such as coconut fatty acids esterified with 20 isethionic acid and neutralized with sodium hydroxide; sodium and ammonium salts of fatty acid amides of methyl taurine; alkane monosulfonates such as those derived by reacting alpha-olefins (C_8-C_{20}) with sodium bisulfite and those derived by reacting paraffins with SO_2 and Cl_2 and 25 then hydrolyzing with a base to produce a random sulfonate; sodium and ammonium $C_7 - C_{12}$ dialkyl sulfosuccinates; and olefinic sulfonates, which term is used to describe the material made by reacting olefins, particularly $C_{10}\text{--}C_{20}$ alpha-olefins, with SO_3 and then neutralizing and 30 hydrolyzing the reaction product. The preferred anionic detergent compounds are sodium $(C_{11}-C_{15})$ alkylbenzene sulfonates; sodium $(C_{16}\text{--}C_{18})$ alkyl sulfates and sodium

35 Examples of suitable nonionic surface-active compounds which may be used preferably together with the anionic surface-active compounds include, in particular, the

 $(C_{16}-C_{18})$ alkyl ether sulfates.

reaction products of alkylene oxides, usually ethylene oxide, with alkyl (C_6-C_{22}) phenols, generally 2-25 EO, i.e. 2-25 units of ethylene oxide per molecule; the condensation products of aliphatic (C_8-C_{18}) primary or secondary linear or branched alcohols with ethylene oxide, generally 2-30 EO, and products made by condensation of ethylene oxide with the reaction products of propylene oxide and ethylenediamine. Other so-called nonionic surface-actives include alkyl polyglycosides, polyhydroxy fatty acid amides (e.g. $C_{12}-C_{18}$ N-methyl glucamide), long-chain tertiary amine oxides, long-chain tertiary phosphine oxides and dialkyl sulfoxides.

Amounts of amphoteric or zwitterionic surface-active

compounds can also be used in the compositions of the invention but this is not normally desired owing to their relatively high cost. If any amphoteric or zwitterionic detergent compounds are used, it is generally in small amounts in compositions based on the much more commonly used synthetic anionic and nonionic actives.

Soaps may also be incorporated into the compositions of the invention, preferably at a level of less than 30% by weight. They are particularly useful at low levels in 25 binary (soap/anionic) or ternary mixtures together with nonionic or mixed synthetic anionic and nonionic compounds. Soaps which are used are preferably the sodium or, less desirably potassium, salts of saturated or unsaturated $C_{10}\text{--}C_{24}$ fatty acids or mixtures thereof. The amount of such 30 soaps can be varied between about 0.5 and about 25% by weight, with lower amounts of about 0.5 to about 5% being generally sufficient for lather control. Amounts of soap between about 2 and about 20%, especially between about 5 and about 15%, are used to give a beneficial effect on 35 detergency. This is particularly valuable in compositions used in hard water when the soap acts as a supplementary builder.

The detergent compositions of the invention will normally also contain a detergency builder. Builder materials may be selected from (1) calcium sequestrant materials; (2) precipitating materials; (3) calcium ion-exchange materials; and (4) mixtures thereof.

In particular, the compositions of the invention may contain any one of the organic or inorganic builder materials, such as sodium or potassium tripolyphosphate, sodium or potassium pyrophosphate, sodium or potassium orthophosphate, sodium carbonate, the sodium salt of nitrilotriacetic acid, sodium citrate, carboxymethylmalonate, carboxymethyloxysuccinate, tartrate mono- and di-succinates, oxydisuccinate, crystalline or amorphous aluminosilicates and mixtures thereof.

Polycarboxylic homo- and copolymers may also be included as builders and to function as powder structurants or processing aids. Particularly preferred are polyacrylic acid (available under the trademark Acrysol from the Rohm and Haas Company) and acrylic-maleic acid copolymers (available under the trademark Sokalan from the BASF Corporation) and alkali metal or other salts thereof.

These builder materials may be present at a level of, for example, from 1 to 80% by weight, preferably from 10 to 60% by weight.

Upon dispersal in a wash water, the initial amount of
peroxygen compound should range in amount to yield anywhere
from about 0.05 to about 250 ppm active oxygen per liter of
water, preferably between about 1 to 50 ppm. Within the
wash media, the amount of oxygen transfer agent initially
present should be from about 0.01 to about 300 ppm,
preferably from about 1 to 100 ppm. Surfactant should be

preferably from about 1 to 100 ppm. Surfactant should be present in the wash water from about 0.05 to 1.0 grams per liter, preferably from 0.15 to 0.20 grams per liter. When

WO 95/13352

present, the builder amount will range from about 0.1 to 3.0 grams per liter.

Apart from the components already mentioned, the detergent 5 compositions of the invention can contain any of the conventional additives in the amounts in which such materials are normally employed in detergent compositions. Examples of these additives include lather boosters such as alkanolamides, particularly the monoethanolamides derived 10 from palmkernel fatty acids and coconut fatty acids, lather depressants such as alkyl phosphates and silicones, antiredeposition agents such as sodium carboxymethylcellulose and alkyl or substituted alkylcellulose ethers, other stabilizers such as 15 ethylenediaminetetraacetic acid, fabric softening agents, inorganic salts such as sodium sulfate and, usually present in very small amounts, fluorescent whitening agents, perfumes, enzymes such as proteases, cellulases, lipases and amylases, germicides and colorants. 20

Stained consumer products benefiting from treatment with compositions of this invention may include clothes and other fabrics; household fixtures and appliances such as sinks, toilet bowls and oven ranges; tableware such as 25 drinking glasses, dishes, cookware and utensils; and even dentures. Hair colorants may also be formulated with the bleach composition of this invention. The bleaching system of this invention may also be applied to industrial uses such as for the bleaching of wood pulp.

30

The system of the present invention may be delivered in a variety of product forms including powders, on sheets or other substrates, in pouches, in tablets, in aqueous liquids, or in non-aqueous liquids such as liquid nonionic 35 detergents.

WO 95/13352

The following examples will more fully illustrate the embodiments of this invention. All parts, percentages and proportions referred to herein and in the appended claims are by weight unless otherwise illustrated.

5

EXAMPLE 1

3,4-Dihydroisoquinoline

The title compound was prepared, using the procedure of Yamazaki, Chem. Lett. (1992) 823. The starting reagent 1,2,3,4-tetrahydroisoquinoline was distilled under vacuum at 85°C (~10 mm Hg) prior to use.

Into a 3-necked 12 L Morton flask equipped with a 10 mechanical stirrer were placed 1,2,3,4tetrahydroisoquinoline (66.6 grams, 0.5 mol) and 2 liters of methylene chloride. A solution of potassium persulfate (189.23 grams, 0.7 mol), sodium hydroxide (48 grams, 1.2 mol) and water (4.4 L) was added. While the solution was 15 vigorously mixed, a 0.05 M aqueous solution of nickel sulfate hexahydrate (200 ml, 0.01 mol) was added to the biphasic mixture. Fine black precipitates deposited immediately, and the resulting mixture was stirred vigorously at room temperature. The reaction temperature 20 became slightly elevated to about 35°C. After 3.5 hours of stirring, the black color changed to a light brown colored solution. The mixture was filtered through a short Celite frit column to remove any insoluble materials. The organic layer in the filtrate was separated by extraction with 25 methylene chloride. The organic layer was dried over magnesium sulfate and concentrated down to give 56 g (85% yield) of a dark red liquid. Distillation at 105°C under vacuum (~11 mm Hg) provided 46 g (70%) of a yellow liquid.

30 The sample contained two impurities: 4% of isoquinoline and 1% of unreacted 1,2,3,4-tetrahydroisoquinoline.

¹H NMR (CDCl₃/TMS): δ 2.73 (t,2H) 3.77 (t,2H), 7.09-7.37 (m,4H), 8.33 (s,1H). The isoquinoline is shown at δ 3.10 (t,2H), 4.01 (s,2H) 7.59-7.94 (m,4H), 8.51 (d,2H) and 9.25 (s,1H).

N-Methyl-3,4-dihydroisoguinolinium p-toluenesulfonate (Imine Quat OTs)

The title compound was prepared, using the procedure described by Koelsch et al. in J. Am. Chem. Soc. (1953) 75, 5 2095. In a 250-ml one-necked round-bottomed flask equipped with a magnetic stir bar and a reflux condenser were placed 3,4-dihydroisoquinoline (8.27 grams, 63 mmol) and 40 ml of methanol. The mixture was stirred and cooled to 0°C. A solution of methyl p-toluenesulfonate (11.73 g, 63 mmol)

10 and 70 ml of methanol was added dropwise. The color of the solution remained clear and yellow. The mixture was heated to reflux for 6 hours. The methanol was then removed in vacuo to provide a quantitative yield (20 g) of an offwhite solid (mp 115-119°C). The solid was pulverized and 15 washed with 40 ml of acetone at room temperature.

Suction filtration provided a white solid in 88% yield: mp

 ^{1}H NMR (D₂O): δ 2.27 (s,3H), 3.15 (t,2H), 3.73 (S,3H), 3.93 20 (t,2H), 7.18-7.78 (m,8H), 8.77 (s,1H). Impurities observed at 4.46 (s), 8.0-8.3 (m), 9.4 (s) were due to N-methylisoquinolinium p-toluenesulfonate.

EXAMPLE 2

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N-Methyl-3,4-dihydroisoquinolinium borontetrafluoride (Imine Quat BF₄)

The title compound was prepared according to the literature procedure (Hanquet, G., Lusinchi X., Milliet, P.,

30 <u>Tetrahedron Letters</u>, (1988), <u>29</u>, 3941).

In a 50 mL 2-necked round-bottomed flask equipped with a reflux condenser and stir bar under nitrogen were placed 3,4-dihydroisoquinoline (1.0 g, 7.6 mmol) and 30 mL of 35 anhydrous toluene. Once in solution, trimethyloxonium tetrafluoroborate (1.12 g, 7.6 mmol) was added, which was not soluble in toluene. The reaction mixture was stirred at WO 95/13352 PCT/EP94/03656

21

room temperature for 10 hours. The reaction mixture was separated into two levels. The dark-red brown viscous liquid was decanted out and dried in an oven to remove any excess toluene.

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¹H NMR (DMSO/TMS): δ 3.23 (2H,m), 3.73 (3H,s), 4.02 (2H,s), 7.47-7.82 (4H,m) and 9.18(1H,s).

EXAMPLE 3

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N-(n-Octyl)-3,4-Dihydroisoguinoline p-toluenesulfonate (Octyl Quat OTs)

3,4-Dihydroisoquinoline (1.31 g, 10.0 mmol) was dissolved in 3 ml MeOH in a 3-neck 25 ml round bottom flask fitted 15 with a condenser, drying tube and stirrer bar, and cooled in an ice bath. n-Octyl p-toluenesulfonate (2.84 g, 10.0 mmol)*, dissolved in 7.0 ml methanol, was added dropwise over about 7 minutes via an addition funnel. The ice bath was removed and replaced with an oil bath and the colorless 20 clear solution heated to reflux for at least 8 hours during which time the reaction solution turned yellow. Removal of the methanol in vacuo gave slightly colored solid product which was triturated with about 7.5 ml acetone. The acetone insoluble solids were filtered, washed with more acetone 25 and dried in a vacuum desiccator. Yield of colorless solids was 1.91 g. A second batch of solids was recovered from the acetone filtrate/washing and filtered, washed with acetone and dried to give 0.31 gm colorless solids. H NMR (CDCl3, 200 MHz) δ 9.70 (1H, s, C₁), 7.00-8.10 (8H, m), 4.20 (2H, 30 t), 4.00 (2H, t), 3.25 (2H, t), 2.30 (3H, s), 1.80 (2H, m), 1.20 (10H, m), 0.87 (3H, t).

*C.S.Marvel and V.C.Seker, <u>Org. Syn.</u>, Coll. Vol. III (1755), p. 366.

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EXAMPLE 4

Stain bleaching experiments were conducted in a Terg-OTometer in 500 mL of milli-Q water, using two tea-stained

5 cotton cloths measuring 3x4 inches. In a typical test, 0.75
g of commercial detergent was added to the system and the
pH of the solution was constantly buffered to the indicated
level by the addition of dilute aqueous sodium hydroxide or
hydrochloric acid. A given oxidant was then added to the

10 system, followed by an appropriate amount of quaternary
imine salt. Washes were carried out at the indicated
temperature for 15 minutes.

Stain bleaching was measured reflectometrically, using a

15 Colorgard System/05 Reflectometer. ΔR is the reflectance difference between washed and unwashed cloths; effects due to detergent are not subtracted. Bleaching was indicated by an increase in reflectance, reported as ΔΔR. In general, a ΔΔR of one unit is perceivable in a paired comparison while 20 ΔΔR of two units is perceivable monadically.

Table II and III report the bleaching activity of the N-methyl-3,4-dihydroisoquinoline borontetrafluoride (BF $_4$) and p-toluenesulfonate (OTs) salts.

TABLE II
Tea Stain (BC-1) at pH 10 and 18°C

BASE* PEROXIDE SALT (10 ppm a.o) COUNTERION P-SURF® OXONE BF4 ULTRA SURF® OXONE BF4					
OXONE OXONE OXONE	SALT	IMINE QUAT SALT CONCENTRATION	AR PEROXIDE	AR IMINE QUAT SALT	AAR IMINE QUAT SALT
OXONE OXONE OXONE				T T ENOUT DE	
OXONE OXONE	BF	6.0X10.5M	200	10.01	
OXONE JRF® OXONE			00:0	#C.01	90.00
OXONE	BF,	M ₉ .01X0.9	1 16		
OXONE			1.10	1.64	6.08
	BF	6.0X10.6M	0	C T	
			00:0	67.0	4.59
ULTRA ALL® OXONE BF4	BF4	6.0X10.6M	700	0	4 ,
			10.0	20.0	5.15

*P-Surf® : Anionic/nonionic/phosphate builder; dosage 1.5 g/l. Ultra-Surf® : 15% anionic/8% nonionic/30% zeolite/20% carbonate; dosage 1.02-1.1 g/l. Ultra-All® : 14% anionic/38% carbonate/29% sulfate; dosage 1.02-1.1 g/l.

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TABLE III

Tea Stain (BC-1) at varying pH and 10°C

	BASE	3ASE PEROXIDE	Нф	SALT COUNTERION	IMINE QUAT	AR IMINE PEROXIDE	AAR IMINE QUAT SALT	AAR IMINE OOR IMINE QUAT SALT QUAT SALT	
					CONCENTRALION		& PEROXIDE		
	ULTRA SURF®	OXONE	10.0	oTs	6,0x10 ⁵ M	1	11.2		
,									
) T	SURF®	PAP*	9.5	OTS	6.0x10 ⁵ M	5.3	13.2	7.9	

*N,N-phthaloylamino-peroxycaproic acid.

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WO 95/13352 PCT/EP94/03656

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Based on the results in Tables II and III, it is evident that the imine quat salts have a pronounced effect upon improving bleaching of a tea-stained cloth. Different formulated base powders have a relatively small effect on the bleaching performance of the imine quat salt.

EXAMPLE 5

This Example illustrates the effect of pH upon a system

10 utilizing the imine quat salt of Example 1 and PAP as the
peroxide source. There is relatively little change with
respect to bleaching performance over the pH range 8.0 to

TABLE IV

Tea-stained Cloth at an 18°C Wash Temperature

рн	IMINE QUAT SALT CONCENTRATION	ΔR PEROXIDE	AR IMINE QUAT SALT & PEROXIDE	ΔΔR IMINE QUAT
8.0	6.0x10 ⁵ M	5.60	14.0	SALT
9.0	6.0x10 ⁵ M	4.90	15-1-27	8.40
10.0	6.0x10 ⁵ M	4.20	13.0	8.10

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EXAMPLE 6

This Example details the effect against stains other than that of tea. Table V establishes that the imine quat salt of Example 2 enhances the bleaching performance of OXONE against a spaghetti sauce stain.

TABLE V

Spaghetti Sauce-stained Cloth at an 18°C Wash Temperature

BASE	PEROXIDE	IMINE QUAT SALT CONCENTRATION	AR PEROXIDE	ΔR IMINE QUAT SALT & PEROXIDE	ΔΔR IMINE QUAT SALT
ULTRA ALL®	OXONE	6.0x10 ⁵ M	13.65	16.53	2.88

Wine (EMPA-114) stains were found to be effectively removed 10 by the imine quat salt of Example 1 in combination with PAP as the peroxide source. Table VI outlines these results.

TABLE VI

Wine-stained Cloth at an 18°C Wash Temperature

BASE	PEROXIDE	IMINE QUAT SALT CONCENTRATIO N	ΔR PEROXIDE	AR IMINE QUAT SALT & PEROXIDE	ΔΔR IMINE QUAT SALT
ULTRA ALL®	PAP	6.0x10 ⁵ M	14.8	18.1	3.3

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EXAMPLE 7

This Example illustrates the performance of the octyl imine quat salt of Example 3 and PAP as the peroxide source. Surprisingly, at lower temperatures, the performance was better than at higher wash temperatures.

TABLE VII

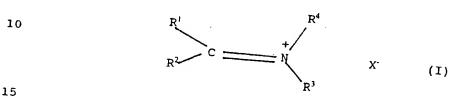
Tea Stain (BC-1) Bleaching Performance of Octvl Quat pH 9.5, 1.02 g/l Ultra All®, 15-minute wash

	TEMP. (0°C)	PAP (7.5 ppm a.o)	ΔR PAP	ΔR PAP +	OCTYL IN	INE QUAT
				6x10 ⁶ M	2x10 ⁵ M	6x10 ⁵ M
- 1	10	Powder in DMF	2.1	6.2	8.3	
	10	Granule	3.7			9.6
	32	Powder in DMF		7.8	10.3	12.1
I	32		4.1	5.6	6.1	6.9
Œ	32	Granule	5.6	6.4	7.4	9.5

The foregoing description and Example: illustrate selected embodiments of the present invention. In light thereof, various modifications will be suggested to one skilled in the art, all of which are "ithin the spirit and purview of this invention.

CLAIMS

- A bleaching composition comprising:
- (i) from about 1 to about 60% by weight of a peroxygen 5 compound;
 - (ii) from about 0.01 to about 10% of an oxygen transfer agent whose structure is:



wherein:

 R^{1} and R^{4} may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl and cycloalkyl radicals;

- R^2 may be a substituted or unsubstituted radical selected 20 from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, cyano, alkoxy, keto, carboxylic and carboalkoxy radicals;
- ${\ensuremath{\mathsf{R}}}^3$ may be a substituted or unsubstituted radical selected 25 from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, and cyano radicals; R^{1} with R^{2} and R^{2} with R^{3} may respectively together form \boldsymbol{a} radical selected from the group consisting of cycloalkyl, polycyclo, heterocyclic and aromatic ring systems;
- X is a counterion stable in the presence of oxidizing 30 agents; and
 - (iii) from about 0.5 to 50% of a surfactant.
- A composition according to claim 1, further comprising from about 1 to about 80% of a detergent

3. A composition according to claim 1, further comprising an effective amount for cleaning of an enzyme selected from the group consisting of proteases, cellulases, lipases, amylases and mixtures thereof.

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- 4. A composition according to claim 1 delivered in a form selected from the group consisting of a powder, sheet, pouch, tablet, aqueous liquid and non-aqueous liquid.
- 10 5. A composition according to claim 1 wherein the peroxygen compound is present in an amount from about 1.5 to 25% and the oxygen transfer agent is present in an amount from about 0.2 to 5% by weight.
- 15 6. A composition according to claim 1 wherein the peroxygen compound is an inorganic material selected from the group consisting of perborate, percarbo te, perphosphate, persilicate and monopersulphate salts.
- 20 7. A composition according to claim 1 wherein the peroxygen compound is an organic peroxyacid.
- 8. A composition according to claim 5 wherein the organic peroxyacid is selected from the group consisting of peracetic acid, monoperoxyphthalic acid, diperoxy dodecanedioic acid, N,N'-terephthaloyl-di(6-aminoperoxy caproic acid) and N,N'-phthaloylaminoperoxycaproic acid.
- A composition according to claim 7 wherein the
 organic peroxyacid is an amidoperoxyacid.
 - 10. A composition according to claim 1 wherein said substituent on R^1 , R^2 , R^3 and R^4 is a functional unit selected from the group consisting of nitro, halo, cyano,
- 35 C_1-C_{20} alkyl, amino, aminoalkyl, thioalkyl, sulfoxyalkyl, carboxyester, hydroxy, C_1-C_{20} alkoxy, polyalkoxy, C_1-C_{40} quaternary di- or tri-alkylammonium functional units and

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mixtures thereof.

- 11. A composition according to claim 1 wherein the oxygen transfer agent is 3,4-dihydroisoquinolinium salt.
- 12. A composition according to claim 11 wherein the salt is an N-methyl-3,4-dihydroisoquinolinium salt.
- 13. A composition according to claim 11 wherein the salt 10 is an N-octyl-3,4-dihydroisoquinolinium salt.
- 14. A method for bleaching a stained substrate, said method comprising contacting said stained substrate in an aqueous medium with a peroxygen compound, a surfactant in an effective amount to clean said substrate and with an oxygen transfer agent whose structure is:

wherein:

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R¹ and R⁴ may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl and cycloalkyl radicals;

R² may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, cyano, alkoxy, keto, carboxylic and carboalkoxy radicals;

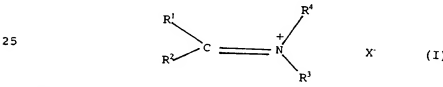
R³ may be a substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, and cyano radicals;

 R^1 with R^2 and R^2 with R^3 may respectively together form a radical selected from the group consisting of cycloalkyl, polycyclo, heterocyclic and aromatic ring systems;

X is a counterion stable in the presence of oxidizing agents; and said peroygen compound to oxygen transfer agent

being present in a molar ratio ranging from about 1500:1 to about 1:2.

- 15. A method according to claim 14 wherein the ratio of peroxygen compound to oxygen transfer agent ranges from about 150:1 to 1:1.
- 16. A method according to claim 14 wherein said substrate is selected from the group consisting of fabrics,10 household fixtures and tableware.
 - 17. A method according to claim 14 wherein said substrate is a denture.
- 15 18. A composition according to claim 14 wherein the oxygen transfer agent is a 3,4-dihydroisoquinolinium salt.
- 19. A method for bleaching a stained substrate, said method comprising contacting said stained substrate in an20 aqueous medium with a peroxygen compound and with an oxygen transfer agent whose structure is:



wherein:

R may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl and cycloalkyl radicals;

R² may be a substituted or unsubstituted radical selected from the group consisting of hydrogen, phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, cyano, alkoxy, keto, carboxylic and carboalkoxy radicals;

R³ may be a substituted or unsubstituted radical and the selections of the selection of

R³ may be a substituted or unsubstituted radical selected from the group consisting of phenyl, aryl, heterocyclic ring, alkyl, cycloalkyl, nitro, halo, and cyano radicals;

 R^1 with R^2 and R^2 with R^3 may respectively together form a radical selected from the group consisting of cycloalkyl, polycyclo, heterocyclic and aromatic ring systems;

- X⁻ is a counterion stable in the presence of oxidizing 5 agents; and said contacting to achieve bleaching being performed in said medium containing about 0.05 to about 250 ppm active oxygen from the peroygen compound and about 0.01 to about 300 ppm oxygen transfer agent.
- 10 20. A method according to claim 19 wherein the oxygen transfer agent is present from about 5 to about 100 ppm per liter of medium.
- 21. A method according to claim 19, further comprising from about 0.05 to about 1.0 grams surfactant per liter of medium.

Inte mal Application No

PCT/EP 94/03656 A. CLASSIFICATION F SUBJECT MATTER IPC 6 C11D3/39 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 6 Clid Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ' Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. US,A,5 041 232 (D.J. BATAL ET AL.) 20 1-7,11, August 1991 14-21 cited in the application see the whole document US,A,4 210 551 (G.R. BRUBAKER ET AL) 1 1,2,4-6, July 1980 14, 16, 19 see claims US,A,4 194 987 (G.R. BRUBAKER ET AL) 25 1,2,4-6, March 1980 see claims FR,A,2 222 428 (AMERICAN CYANAMID) 18 1,14 October 1974 see claims; examples -/--X Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documents: T later document published after the international filling date or priority date and not in conflict with the application but died to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date 'L' document which may throw doubts on priority claim(s) or which is cited to enablish the publication date of another citation or other special reason (as specified) Y document of particular relevance; the claimed invention camot be conndered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "O" document referring to an oral disclosure, use, exhibition or Other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 17. 03. 95 13 March 1995 Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2230 HV Rujewijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Authorized officer

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INTERNATIONAL SEARCH REPORT

Inte mal Application No PCT/EP 94/03656

Category *	DOOD DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/EP 94/03656	
gury	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	_
A	DATABASE WPI Derwent Publications Ltd., London, GB:	1,14	_
	& JP,A,2 173 098 (LION CORP) 4 July 1990 see abstract		

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INTERNATIONAL SEARCH REPORT

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Inter cal Application No

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Patent documen: cited in search report	Publication date	Pat	ent family	P 94/03656
US-A-5041232	20-08-91	m	ember(s)	Publication date
	50-00-31	AU-B- AU-A- CA-A- DE-D- DE-T- EP-A- ES-T- JP-A-	7287001	06-05-93 19-09-91 17-09-91 10-11-94 09-02-95 23-10-91 01-12-94
US-A-4210551	01-07-80	NONE		17-08-92
US-A-4194987	25-03-80	NONE		
FR-A-2222428	18-10-74	US-A- AU-A- BE-A- CA-A- DE-A- GB-A- JP-A- NL-A-	3824188 6571974 812561 1022304 2412954 1424502 49126580 7403434	16-07-74 21-08-75 20-09-74 13-12-77 26-09-74 11-02-76 04-12-74 24-09-74

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